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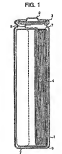
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- (54) ELECTROLYTE FOR NON-AQUEOUS CELL AND NON-AQUEOUS SECONDARY CELL
- (57) In a rechargeable non-aqueous electrolyte secondary battery using positive electrodes, negative electrodes and a non-aqueous electrolytic solution, additives to the electrolytic solution are used in combination. preferably in combination of at least two compounds selected from o-terphenyl, triphenylene, cyclohexylbenzene and biphenyl, and thus there are provided batteries excellent in safety and storage characteristics.



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## Description

#### TECHNICAL FIELD

5 [0001] The present invention relates to a non-aquious electrolytic solution which can ensure safety of batteries at the time of overcharging with improving recovery characteristics of the batterios after storage at high temperatures, and to a non-aqueous type secondary battery using said electrolytic solution.

## BACKGROUND ART

- [0002] Rocently, AV devices and electronic devices such as porsonal computers of portable or corcless type have been rapidly developed, and secondary batteries which are arreall in size, light in weight and high in energy density are sementally demanded as electric sources for chirty these devices. Among them, non-equeues electroity is solution secondary batteries using a negative electroit containing thirum as an active material are hopefully expected as satteries having high voltage and high energy density.
- [0003] In the above batteries, lithium-containing metal oxides which show a voltage on the order of 4 V are used for positive electrode active materials, and materials capable of intercalation or delintercalation of lithium, such as carbon-acous materials, are used for neartive electrories.
- [0004]. One of the most important tasks in these non-aqueous electrolytic solution batteries is to ensure the safety.

  [0005] Particularly, in lithium ion secondary batteries, when they are charged in excess of a given charging voltage
  due to, for example, troubles of charging control dirouts, they are in overcharged each, and lithium nos in the possive
  electrode are excessively extracted and migrate to negative electrode to eause absorption of lithium in an amount
  larger than the prescribed design capacity in the negative electrode to re causes predipitation of lithium as metallic
  lithium on the surface of negative electrode. It the batteries in euch a state are further forcedly charged, internal missteance of the batteries increases and generation of head due to the Joule's heat becomes great to cause ahormant heat
- an ance of the batteries increases and generation of heat due to the Joule's heat becomes great to cause abnormal heat generation, and, in the worst case, to result in hemmal runaway. By providing a current interrupting switch of temperature sensing type (for example, a positive temperature occellicient themsitor (PTC) or a temperature fuse) outside the batteries, the current is interrupted without fall, and safety can be ensured at the time of generation of abnormal heat. The providing outside the providing of internal pressure of batteries is generally employed as disclosed in U.S. Palant No. 4943497.
- [0006] However, in the case of using such a mechanical current interrupting mechanism, reduction does it difficult and, furthermore, with the batteries becoming smaller and thinner, it becomes structurally difficult to insert the mechanism in the horarise.
- [0007] For solving the above problems, there is proposed a method of adding to the electrolytic solution an additive as which causes a reversible redox reaction, thereby to self-consume the electric energy introduced into the batteries as a radox shittle (for example, PA-4-2968T, IV-PA-9308T, etc.).
  - [0008] However, in the method of using the redox shuttle, when the overcharging current becomes great, there are limits in charge transfer reaction rate and lithium ion transfer rate, and, thus, it cannot be said that the method provides a sufficient safety.
- 40 [0009] For solving the above problems, JP-9-50822, JP-A-10-50342, JP-0-106355, JP-10-21256, Jupenace Patent No. 2939-969, and JP-A-2000-58117 propose a means of adding to batteries an aromatic compound having a methopy group and a halogen group, bipheny for thiophene, or an aromatic other compound, which polymerizes at the time of overcharging to result in fining of termorestime and, bus, to ensure the safety.

## 45 DISCLOSURE OF INVENTION

- [0010] In the case of the batteries in which a current interrupting switch of temperature sensing type (for example, a positive temperature coefficient thermistor (PTC) or a temperature (use) is provided outside the batteries against abnormal heat generation, when an excessively large overcharging current (5-6 C) of more than 5-8 times the rated 20 capacity passes, the device per se generates heaf due to the current and resistance increases, thereby interrupting the current, and thus the selfey can be ensured, but in the case of a current generally used for charging and discharging the batteries (tess than 1-2 C at the maximum), the rising of temperature is insufficient and the resistance does not increase. When overcharging is carried out at such a current value, the safety cannot be sufficiently ensured. If the safety is the carried of the harderies are demands.
  - [0011] In the case of adding to the batteries the above-mentioned additives such as aromatic compound having a methoxy group and a halogen group, highenyl or thiophene, and an aromatic either compound, it has been confirmed that in the generally used current ranges, the additives polymerize on the electrodes in overhanged state, and the

safety is improved.

[0012] However, it has been found that these additives must be added in an amount of not less than 1% by weight for ensuring the safety at the time of overcharging, but if the additives are added in a large amount, in an shell life test, for example, an environment set (60°C) which supposes the case of leaving them in a car in summer, these additives partially react to cover the active material, resulting in considerable deterioration of the battery characteristics.

[0013] The above phenomenon is considered to occur because oxidative polymenization potential of the additives lowers due to exposure of the battery to a high temperature environment, and, furthermore, the potential distribution in the charged electrode is not uniform and higher potential profits are partially present, and, as a result, the additives react even in the ordinary environment of potential at which batters is used.

[0014] The above problem of deterioration due to storage can be solved, for example, by using an additive high in oxidative polymerization starting potential (for example, cyclinderybcancae), but in this case, since the reaction to tail and ot the additive by the companies of the overcharging cannot sufficiently be ensured.

[0015] The present invention solves the above problems and to provide a battery excellent in high-temperature storage characteristics while ensuring the safety at overcharging.

[0016] In order to solve the above problems, according to the present invention, in a non-aqueous electrolytic solution In which an electrolyte is dissolved in a non-aqueous solvent, two or more organic compounds differing in exidative polymerization reaction potential are added. Preferably, organic compounds of relatively low exidation reaction potential are added in a very small amount, preferably not less than 0.01% by weight and less than 1.0% by weight based on

2b the total amount of the electrolytic solition, thereby to control the recovery characteristics are strongly as the total amount of the electrolytic solition, thereby to control the recovery characteristics after storage and the safety during overcharging. Specifically, it is pretend to add at least two organic compounds selected from o-terphenyl, triphenylene, cytólchezylenzare and biphonyl.

[0017] It is preferred that not less than 1.0% by weight and not more than 3.0% by weight of o-terphenyl and not less than 0.01% by weight and less than 1.0% by weight of triphenylene are contained in the non-equieous solvent.

25 [0018] Furthermore, it is preferred that not less than 1.0% by weight and not more than 5.0% by weight of cyclohexylbenzene and not less than 0.01% by weight and less than 1.0% by weight of biphanyl are contained in the nonaqueous solvent.

[0019] Moreover, it is preferred that not less than 1.0% by weight and not more than 5.0% by weight of cyclohexylbenzene and not less than 0.01% by weight and less than 1.0% by weight of o-temphenyl are contained in the nonaqueous solvent.

[0000] Further, it is preferred that not less than 1.0% by weight and not more than 5.0% by weight of cyclohexylbencene, not less than 0.01% by weight and less than 1.0% by weight of o-terphenyl and not less than 0.01% by weight and less than 1.0% by weight of biphenyl are contained in the non-equous solvent.

[0021] Furthermore, it is preferred that all of o-tephenyl, triphenylere, cyclohazy/benzene and biphenylare contained in the non-auguous solvent and the total amount of them to 0.4-5% by weight based on the non-auguous solvent. [0022] These organic compounds are particularly effective when the positive electrodes comprise a material containing a lithium-containing metal oxide and the negative electrodes comprise a material containing graphite, and the non-auguous electrofytic solution exerts the higher effect when it is an electrofytic solution in which all think metal as a

solute la dissolved in a non-aqueous solvent mainly composed of a cyclic carbonate and a chain carbonate.

[1023] The cyclic carbonate is preferably at teast one compound selected from ethylene carbonate (EC), propylone carbonate (PC), butylene carbonate (PC), butylene carbonate (PC).

[0024] The chain carbonate is preferably at least one compound selected from dimethyl carbonate (DMC), diethyl carbonate (DEC), ethylmethyl carbonate (EMC), methylpropyl carbonate (MPC) and ethylpropyl carbonate (EPC).

## 45 BRIEF DESCRIPTION OF DRAWING

#### [0025]

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FIG. 1 is a longitudinal sectional view of a cylindrical battery in the examples of the present invention and in the comparative examples.

# BEST MODE FOR CARRYING OUT THE INVENTION

[0026] In the present invention, recovery characteristics efter storage and safety during overcharging can be controlled by adding two or more organic compounds differing in oxidative polymerization reaction potential to the electrolitic solution.

[0027] Examples of organic compounds differing in exidative polymerization potential contained in the electrolytic solution for non-aqueous type batteries in which an electrolyte is dissolved in a non-aqueous solvent are o-temperyl,

tiphenylmer, cyclohoxybenzene and biphenyl. As to the content of the organic compounds, the weight of the organic compounds of relatively higher coldidative polymerization potential is preferably not less than 1.0% by weight and not more has 6.0% by weight based on the total amount of the non-aqueous electrolytic solution. The veight of the organic compound of the relatively lower coldistive polymerization potential is prierably not less than 0.0% by weight hased on the total amount of the non-aqueous electrolytic solution. Furthermore, the weight ratio than 1.0% by weight based on the total amount of the non-aqueous electrolytic solution. Furthermore, the weight ratio the organic compound of relatively higher codative popherization reaction potential and the organic compound of relatively lower oxidative polymerization reaction potential is preferably not lower than 20:1 and not higher than 2:1, more preferably not lower than 20:1 and not higher than 2:1.

[0028] in order to improve the recovery-theractoristics after storage, the amount of the organic compound of relatively lower oxidiative polymerzation starting potential (for example, bipheraply) is perfeately smaller, but in order to ensure the safety at overcharging, the organic compound must react as much as possible at overcharging, namely, the amount is preferrably rather largor.

[0029] According to the present invention, two or more organic compounds differing in oxidative polymerization reaction potential (herelinates rosentieme relevant to as "additives") are used, and the amount of the organic compound
for its active polymerization starting potential (for example, bisheny) is conspicuously reduced in this
system, thereby maintaining excellent storage characteristics, and on the other hand the organic compounds resart
only slightly at the overcharging, whereby polarization at the overcharging increases, and the organic compound or
relatively higher oxidative polymerization starting potential (for example, cyclohexyberzene) resart at an early stage,
and thus the salely can be ensured. As a result, there can be attained both the recovery properties after storage and
the insurance of salely at overcharging, which cannot be attained with addition of each organic compound singly.

one insulative of series at event energing, when cannot be attained with an action or each organic compound singly.

[0030] Since the additives in the present invention do not aim at an action are redox shuttles, the oxidation reaction is desirably interventible and they differ in purpose from JP-A-7-302614 and JP-A-9-50822 which aim at reversibility of redox reaction.

[0031] As the liblum-containing composite oxides used as positive electrode active missterials in the present invention, mention may be made of, for example, ILCO<sub>2</sub>, ILMO<sub>2</sub>, ILMO<sub>3</sub>, ILMO<sub>3</sub>

90 [0032] The lithium-containing composite oxides used as positive electrode active materials in the present invention can be prepared by mixing carbonate, nitrate, oxide or hydroxide of lithium with carbonate, nitrate, oxide or hydroxide of a transition material such as cobalt, mangainese or nickel at a desired composition, grinding the mixture and fifting the powder or by a solution reaction. The fifting method is especially preferred, and the fifting temperature can be 250-1500°C at which a part of the mixed compound is decomposed and motter. The fifting time is preferrably 1-90 shours. The fifting sample sample samp

[0033] In the present invention, a plurality of different positive electrode active materials may be used in combination. [0034] As current collectors of positive electrodes, there may be used any electron conductors as long as they do not undergo chemical changes in the constructed batteries. For examples, as materials of the current collectors, there may be used stainless steel, aluminum, tilanium and carbon, and aluminum or aluminum alloys are especially preferred. As for the shape of the current collectors, they may be in the form of foil, film, sheet, net, punched material, lath, porous material, foamed material, fiber group, shaped nonwoven fabric, and the like. The surface of the current collectors may be made rough by a surface treatment, Thickness thereof is not particularly limited, and those of 1-500 um are used. [0035] The negative electrode materials used in the present invention may be lithium alloys, alloys, intermetallic compounds, carbons, organic compounds, Inorganic compounds, metal complexes and organic high molecular compounds, which are capable of absorbing and releasing lithium ions. These may be used each alone or in combination. [0036] As the carbonaceous materials, mention may be made of, for example, cokes, pyrolytic carbons, natural graphite, artificial graphite, mesocarbon microbeads, graphitized mesophase spherules, vapor deposited carbons, glassy carbons, carbon fibers (polyacrylonitrile fibers, pitch fibers, cellulose fibers and vapor deposited carbon fibers). amorphous carbons, and carbons prepared by firing organic materials. These may be used each alone or in combination. Among them, preferred are graphite materials such as those obtained by graphitizing mesophase spherules, natural graphite and artificial graphite. These negative electrode materials may be used as composites, and, for example, combinations of carbon with alloys, carbon with inorganic compounds, and the like can be considered.

[1037] In the present invention, since Lis contained in the positive electrode active material, negative electrode materials which do not contain Li (such as carbon) can be used. Moreover, when Lis added to such negative electrode materials which do not contain Li in a small amount (about 0.01-10 parts by weight based on 100 parts by weight of the negative electrode materials), even if the materials become inactive owing to the reaction of a part of Li with electroty. Li can be supplemented with Li contained in the negative electrode materials, which is preferred, Li can be

contained in the negative electrode materials, for example, in the following manner. That is, lithium metal which is mothen by heating is coated on a current collector to which a negative electrode material is pressed, thereby impregnating the negative electrode material is pressed, thereby impregnating the negative electrode group by press bonding and LI is electrochemically doped in the negative electrode material in the electrody is solution.

[0038] As current collectors of negative electrodes, there may be used any electron conductors as long as they do not undergo chemical changes in the constructed ballories. For example, as malorials of the collectors, there may be used stainless atoel, nickel, copper, filanium, etc., copper or copper adolysa are especially preferred.

[0039] As for the shape of the current collectors, they may be in the form of foil, film, sheet, net, punched material, listh, porus material, former group, shaped nonwoven fabric, and the like. Moreover, the surface of the current collections may be made rough by a surface retainent. This choices is not particularly initiated, and those of 1-500

19040] The non-equipous electrolytic solution in the present invention comprises a solvent and a lithium sait dissolved in the solvent. As the non-equipous solvents, mention in regide to regide controllates used as earlyiers carbonates (EC), proplene carbonates (EC), butylene carbonates (EC), proplene carbonates (EC), butylene carbonates (EC), and whytene carbonates (EC), electrollates such as dimetry carbonate (EC), and the dipropulation (EC), electrollates such as dimetry carbonate (EC), and electrollates (EC), and electrollat

[0041] The Illhium salts which are dissolved in these solvents include, for example, LICIO<sub>4</sub>, LIBF<sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>4</sub>, LIBF<sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>4</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>5</sub>, LIPF<sub>5</sub>, LIPF<sub>5</sub>, LIAIO<sup>1</sup><sub>5</sub>, LIPF<sub>5</sub>, LIPF<sub>5</sub>

[0042] Especially preferable non-squeous electrolytic solution in the present invention is one which contains at least eithylene carbonale and ethylenetyl carbonate and LIFF<sub>6</sub> as a likim salt. The armount of the electrolytic solution contained in the battery is not particularly limited, and it can be used in a necessary amount depending on the amount of positive electrode active material and that on opagive electrode material and that on opagive electrode material and that of nogagive electrode material and that of the battery. The amount of the illustration of the control of th

[0043] The electrolytic solution is ordinarily used by impregnating or filling a separator such as of porous polymer or nonwoven fabric with the electrolytic solution.

[0044] Moreover, there may be used a gelled electrolyte comprising an organic solid electrolyte containing the nonequeuou electrolyte solution. As the organic solid electrolyte, polymente matrix materials such as polyetrylene coxide, polypropytene coxide, polyvinylened flooride and derivatives, mixtures and composites of these materials are effective. Especially preferred are copolymens of vinylidene fluoride and hexafluoropropytene and mixtures of polyvinylidene fluoride and polyteriviene coxide.

[0045] As the separator, an insulating microporous thin film having a high ion permeability and a dealred mechanical strength is used. The separator preferably has a function of closing the pores at a temperature of 80°C or higher to enhance the resistance. Sheets or nonwoven fabrics made from olden polymers comprising one or combination of polyproplene and polyethylene or glass fibers are used from the points of organic solvent resistance and hydrophobic properties. Pore climater of the separator is preferably in such a range that active materials, binders and conducting agants which are released from the electrode sheets do not permeate through the pores, and, for example, the pore climater is preferably 0.01-1 µm. The thickness of the separator is generally 5-300 µm. The porceily is determined depending on the permeability to electron or lost, kind of materials or film thickness, and is desirably 3-0.04.

[0046]. The shape of batteries can be any of sheet type, cylinder type, flat type, rectangular type, etc. When the shape of batteries is sheet type, cylinder type or rectangular type, the mix of positive electrode active material or negative electrode material is used mainly by ocating on a current coffector, then dyring and compressing the collector.

[0047] The shape of the rolled electrodes in the present invention is not necessarily in the form of true cylinder, and may be in the form of ellipsociety cylinder having a ellipsociatel section or in the form of square pillar such as rectangle. [0048] Preferred combinations of interest invention are combinations of the preferred chamical materials and the preferred battery constituting parts mentioned above. Especially preferred are those which contain Li<sub>2</sub>Co<sub>2</sub>, Li<sub>3</sub>Ni<sub>2</sub>O<sub>3</sub>, (Li<sub>3</sub>Xi<sub>2</sub>O<sub>3</sub>) as positive electrode active materials, and acceptance black as a conducting agent. The current collector of positive electrode is made of stainless steel or aluminum, and is in the form of net, sheet, foil or lath. The negative electrode material referred by contains at less one compound such as alloy and carbonoscove material. The

current collector of negative electrode is made of stainless steel or copper and is in the form of net, aheat, foil or lath. Carbon materials such as accepture black and graphite as the electron conducting agent may be contained in here was such as polymers and the properties of th

## EXAMPLES

[0049] Examples of the present invention will be explained below referring to the drawing.

## Example 1

- 20 [0050] FIG. 1 is a longluidinal sectional view of the cylindrical battery used in this example. In FIG. 1, the reference numeral 1 indicates a battery case made by working a stainless steel plate having resistance to organic electrotycic solution, 2 indicates a sealing plate provided with a safety valve, 3 indicates an insulation packing, 4 indicates an electrode plate group, and positive veloctrode valve electrode with separation interposed between the positive electrode and the negative electrode and the positive electrode and the negative electrode and connected to the sealing plate 2, and a negative electrode lead 6 is drawn from the positive electrode and connected to the bottom of the battery case 1. The reference numeral 7 indicates an insulation ring, which is provided at the upper and lower portions of the electrode plate group 4. The positive electrode, the negative electrode, and others will be explained in cleat below.
- [0051] The positive electrode was made in the following manner, Li<sub>2</sub>Co<sub>2</sub> and Co<sub>5</sub>O<sub>4</sub> were mixed and fired at 900°C for 10 hours to prepare an LiCo<sub>2</sub> powder. This powder was mixed with 3% of acetylene black and 7% of a flower on polymer binder based on the weight of the LiCo<sub>2</sub> powder, followed by suspending the mixture in an auguous condownself of the prepared positive electrode mix pasts. The resulting positive electrode mix pasts was coased on the surface of an aliminum foll of 20 min in thickness which was a positive electrode place and the coat was dried, followed by rolling to make a positive electrode place of 0.18 mm in thickness, 37 mm in width and 39 mm in length.
- [0052] For the negative electrode, a mesophase ephanule which was graphitized at a high temperature of 2800°C (hereinalter referred to as "mesophase graphid") was used. This mesophase graphide was mixed with 3% of a sympen-butsdriene rubber based on the weight of the mesophase graphide, and then the returne was suspended in an aqueous carboxymetryl cellulose solvidine to prepare a passet. This negative electrode mixe passet was costed on both aidea of a cost of 10 cl of 0 (0.02 mm in thickness and dried, followed by rolling to make a negative electrode piete of 0.20 mm in thickness, 30 mm in with and 420 mm in length.
- [0053] A lead made of aluminum was attached to the positive electrode plate and a lead made of nickol was attached to the negative electrode plate; and the positive electrode plate end the negative electrode plate with a polyethylene separator of 0.018 mm in thickness, 45 mm in width and 840 mm in length interposed between the positive electrode plate were rolled into a spiral form and it sected in a battery case of 17.0 mm in diameter and 50.0 mm in height. The electrodylic solution used was repeated by dissolving in molifier of LIPE, in anxious observed comprising EC and EMC at a volume ratio of 30.70, and as additives, 2% by weight of o-terphenyl and 0.2% by weight of triphenymen based on the loat amount of the electrolytics obtained may expend to the electrolytics obtained was poured into the battery case, and then the case was sealed to make a battery 1 (battery capacity: 50 mm) of the present invention.

# Example 2

[0054] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that cyclonexylbenzene in an amount of 2% by weight and bipheny in an amount of 2.2% by weight based on the total amount of the electrolytic solution were used as the additives to the electrolytic solution. The thus obtained battery was referred to as battery 2 of the present invention.

## Example 3

[0055] A cylindrical battery of spiral type was mede in the same manner as in Example 1, except that cyclonexylbenzene in an amount of 2% by weight and o-terphenyl in an amount of 0.2% by weight based on the total amount of the electrolytic solution were used as the additives to the electrolytic solution. The thus obtained battery was referred to as battery 3 of the present invention.

## Example 4

10 [0056] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that cyclohexyl-borzene in an amount of 2% by weight, biphonyl in an amount of 2% by weight and o-terphenyl in an amount of 0.2% by weight based on the total amount of the olectrolytic solution were used as the additives to the electrolytic solution. The thus obtained battery was referred to as battery 4 of the present invention.

## 15 Example 5

[0057] A cylindrical battery of spiral type was made in the same manner as in Example 1. except that cyclohaxylbenzene in an emount of 2% by weight, byhenyl in an amount of 0.2% by weight, o-terphenyl in an amount of 0.2% by weight and tyhenylene in an amount of 0.1% by weight based on the total amount of the electricytic solution were used as the additives to the electrolytic solution. The thus obtained battery was referred to as battery's of the present invention.

# Comparative Example 1

25 [0058] A cylindrical battery was made in the same manner as in Example 1, except that the additives to the electrolytic solution were not used. The thus obtained battery was referred to as a comparative battery (battery 6).

## Comparative Example 2

30 [0059] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that biphenyi was used in an amount of 2.0% by weight based on the total amount of 10 the electrolytic solution as the additive to the electrolytic solution. The thus obtained battery was referred to as a comparative battery thattery 7).

# Comparative Example 3

[0060] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that cyclohexylbenzene was used in an amount of 2.0% by weight based on the total amount of the electrolytic soution as the additive to the electrolytic soution. The third sobtained battery was referred to as a comparative battery (battery 5).

## 40 Comparative Example 4

[0061] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that o-terphenyl was used in an amount of 2.0% by weight based on the folial amount of the electrolytic solution as the additive to the electrolytic solution. The thus obtained battery was referred to as a comparative battery (battery 0.3)

#### Comparative Example 5

[0062] A cylindrical battery of spiral type was made in the same manner as in Example 1, except that biphenyl was used in an amount of 0.2% by weight based on the total amount of the electrolytic solution as the additive to the electrolytic solution. The thus obtained battery was referred to as a comparative battery (battery 101).

[0063] Then, as overcharging tests, 20 cells each of the batteries 1-5 of the present invention and the comparative batteries 6-9 were subjected to overcharging at 0.8 A (1C) from charged state at an environmental temperature of 20°C; and it was examined whether shonomal heat generation occurred or not in the batteries. The number of cells among 20 cells in which shonomal heat generation occurred is shown in Tables 1 and 2.

[0084] Furthermore, as high-temperature storage tests, the batteries in changed state were left to stand at 60°C for 4 days, and the discharge capacity at 10 was compared with the discharge capacity before storing, and cepacity recovery rate after storing was calculated. (Capacity recovery rate after storing = (1C discharge capacity before storing) x100(S).

#### Table 1

		IdDIE 1	
L		(Example)	
No.	Additives	Occurrence of abnormal heat generation	Recovery rate after storage (%)
1	o-Terphenyl (2%) + triphenylene (0.2%)	0/20	75
2	Cyclohexylbenzene (2%) + biphenyl (0.2%)	0/20	85
3	Cyclohexylbenzene (2%) + o- terphenyl (0.2%)	2/20	82
4	Cyclohexylbenzene (2%) + biphenyl (0.2%) + o-terphenyl (0.2%)	0/20	84
5	Cyclohexylbenzene (2%) + biphenyl (0.2%) + o-terphenyl (0.2%) + triphenylene (0.1%)	0/20	83

# Table 2

			laule 2				
25	(Comparative Example)						
	No.	Additives	Occurrence of abnormal heat generation	Recovery rate after storage (%)			
	6	No	20/20	87			
	7	Biphenyl (2%)	0/20	17			
30	8	Cyclohexylbenzene (2%)	13/20	85			
	9	o-Terphenyl (2%)	7/20	69			
	10	Biphenyl (0.2)	14/20	82			

- 5 [0065] As shown in Table 2, in the blattery 6 having no additives, the phenomenon of abnormal heat generation occurred in all of 20 cells when they were subjected to overcharging. Hithero, the safety of betteries has been ensured by providing a plurality of safety mechanisms comprising a protective clicuit for charging votage and a current interrupting mechanism, but the abnormal heat generation may occur in case no safety protection was provided as in the Comparative Example 1.
- g0065f) it can be seen that in the battery 7 of Comparative Example 2 in which bipherry was added as the additive, abnormal heat generation did not occur even when it was subjected to overcharging, and safety could be ensured, but the recovery rate after storing at high temporatures was very low, namely, 17%.
- [0067] This bettery was disassembled after storing and analyzed to find a film which was considered to be a polymerization product was formed on the surface of the positive electrode, and it was presumed that the recovery rate decreased due to the hindrance to charing and discharging reaction of lithium ion.
- [0068] Furthermore, in the battery 8 of Comparative Example 3 in which only cyclohoxy/benzene high in reaction starting potential was contained, the recovery rate after storing at high temperatures was high, namely, 85%, and this battery showed characteristics substantially equal to those of the battery 6 containing no additives, but was insufficient in safety at the time of overcharging and more than half of the cells resulted in abnormal heat generation.
- 0 [0069] In the case of addition of o-terphenyl which was relatively high in reaction starting potential, ignition rate at overcharging somewhat decreased, but the decrease was still insufficient and, further, the recovery rate after storing was also insufficient.
  - [0070] When the amount of biphenyl of low reaction starting potential was reduced, the recovery rate after storing increased, but safety at overcharging could not be ensured as in the battery 10 of Comparative Example 5.
- [0071] As explained above, it is not easy to ensure both the safety at overcharging and the high-temperature storage characteristics by using a single additive.
  - [0072] In comparison with the batteries of the comparative examples, in the batteries such as batteries 1-5 of the

present invention in which two or more additives were contained, by adding a small amount of an additive of relatively lower reaction starting potential (bipheny), o'elaphenyl, triphenylene) in each system, the recovery rate after storing was improved and a recovery rate of more than 70% was ministained, and, morrower, since the additive of lower reaction starting potential which slightly reacted during the overcharging increased the potarization at overcharging, the additive of higher reaction starting potential started the reaction at the early stage to enhance the safety at overcharging. Thus, ballowers supplied in recovery characteristics in scoring and high in safety at overcharging could be realized. [0737] From the point of maintaining the safety at overcharging, the secure of contracting the safety at overcharging the safety at over

less than 1.0% by weight and not more than 5.0% by weight.
[0074] Furthermore, from the point of maintaining the recovery characteristics in storing, the amount of biphenyl or
triphenylene is preferably not less than 0.0

## INDUSTRIAL APPLICABILITY

[0075] As expisited above, the present invention can provide batteries having high safety against overcharging and excellent in recovery characteristics in storing at high temperatures by combining additives to electrolytic solutions. [0076] Portable tielephones, portable information terminal devices, carn coders, personal computers, POA, portable audio devices, electric cars, electric sources for road leveling, and the like which are high in safety can be provided by using the non-raqueous type electrolyte secondary batteries as mentioned above.

## Claims

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- A non-aqueous electrolytic solution for non-aqueous type batteries in which an electrolyte is dissolved in a non-aqueous solvent, characterized by containing two or more organic compounds differing in oxidative polymerization potential, with a proviso that weight of the organic compound of relatively lower oxidative polymerization potential is less than that of the organic compound of relatively higher oxidative polymerization potential.
- An electrolytic solution for non-equeous type batteries according to claim 1, wherein two or more organic compounds selected from o-terphenyl, triphenylene, cyclohexylbenzene and biphenyl are contained as the organic compounds differing in oxidative polymerization potential.
  - 3. An electrolytic solution for non-aqueous type batteries according to claim 1, wherein not less than 1.0% by weight and not more than 3.0% by weight of -lerphenyl and not less than 0.01% by weight and less than 1.0% by weight of triphenylene based on the total amount of the non-aqueous electrolytic solution are contained.
- 4. An electrolytic solution for non-equeous type batteries according to claim 1, wherein not less than 1.0% by weight and not more than 5.0% by weight and eless than 1.0% by weight and less than 1.0% by weight of biphonyl based on the total amount of the non-equeous electrolytic solution are contained.
- 49 5. An electrolytic solution for non-equeous type batteries according to claim 1, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohesy/benzene and not less than 0.01% by weight and less than 1.0% by weight of ot-terphenyl based on the total amount of the non-equeous electrolytic solution are contained.
- 6. An electrolytic solution for non-aqueous type batteries according to claim 1, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohexybenzene, not loss than 0.01% by weight and less than 1.0% by weight of c-terphenyl and not less than 0.01% by weight and less than 1.0% by weight of biphenyl based on the total amount of the non-aqueous electrolytic solution are contained.
  - An electrolytic solution for non-aqueous type batteries according to claim 1, wherein all of o-terphenyl, triphenylene,
    cyclohexylberczene and biphenyl are contained and the total amount of thorn is 0.4-5% by weight based on the
    total amount of the non-aqueous electrolitic solution.
  - 8. An electrolytic solution for non-squeous type batteries having a positive electrode comprising a material containing all thism-containing metal oxide as a positive electrode active motorial and an appaive electrode comprising a material containing graphite eas a negative electrode active material, characterized in that the non-equeous electrode in the properties of a cyclic cathonial end as chain cathonials in which is officially endead to a cyclic cathonial end as chain cathonials in which a lithium salt is dissolved as a solute, and further contains two or more organic compounds differting in coldaffve polymerization potential, with a provise that weight of the organic compound of relatively before oxidative.

polymerization potential is less than that of the organic compound of relatively higher oxidative polymerization potential.

- An electrolytic solution for non-aqueous type batteries according to claim 8, wherein two or more organic compounds selected from o-terphenyl, triphenylene, cyclohezy/benzene and biphenyl are contained as the organic compounds differing in oxidative polymerization potential.
- 10. An electrolytic solution for non-equeous type batteries according to claim 8, wherein not less than 1.0% by weight and not more than 3.0% by weight of -terphenyl and not less than 0.01% by weight and less than 1.0% by weight of triphenylenc based on the total amount of the non-equeous electrolytic solution are contained.
- 11. An obstrolytic solution for non-equocus type batterios according to claim 8, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohoxylbenzene and not less than 0.01% by weight and less than 1.0% by weight of biphonyl based on the total amount of the non-equocus electrybic solution are contained.
- 12. An electrolytic solution for non-equeous type batteries according to claim 8, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohaxylbenzene and not less than 0.01% by weight and less than 1.0% by weight of oterpharyl based on the total amount of the non-aqueous electrolytic solution are contained.
- 20 13. An electrolytic solution for non-suapous type batteries according to claim 8, wherein not less than 1.0% by weight, and not more than 5.0% by weight and otherselves there are not less than 0.0% by weight and is sait ban 1.0% by weight and is sait ban 1.0% by weight and the less than 0.0% by weight and the sait ban 1.0% by weight of biphenyl based on the total amount of the non-acqueus electrolytic solution are contained.
- 25 14. An electrolytic solution for non-aqueous type batteries according to claim 8, wherein all of o-terphenyl, triphenylene, cyclohoxyleneane and biphonyl are contained and the total amount of them is 0.4-5% by weight based on the total amount of the non-aqueous electrolytic solution.
- 15. An electrolytic solution for non-aqueous type batteries according to claim 8, wherein the cyclic carbonate is at least so one compound selected from ethylene carbonate (EC), propylene carbonate (PC), butylene carbonate (BC) and vinylene carbonate (VC).
  - 16. An electrolytic solution for non-aqueous type batteries according to claim 6, wherein the chain carbonate is at least one compound selected from dimethyl carbonate (DMC), cliethyl carbonate (DEC), ethylmethyl carbonate (EMC), methyloropyl carbonate (MPC) and ethyloropyl carbonate (FEC).
  - 17. A non-equeous type secondary battary containing a non-aqueous electrolytic solution in which an electrolytic is dissolved in a non-aqueous solvent, characterized in that two or more organic compounds differing in oxidative polymerization potential are lutrier contained in the non-aqueous electrolytic solution, with a provise that weight of the organic compound of relatively lower oxidative polymerization potential less than that of the organic compound of relatively lower oxidative polymerization potential.
- 18. A non-aqueous type secondary battery according to claim 17, wherein two or more organic compounds selected from o-terphenyl, triphenylene, cyclohexybenzene and biphenyl are contained as the organic compounds differing in oxidative polymerization potential.
  - 19. A non-equeous type secondary battery according to claim 17, wherein not less than 1.0% by weight and not more than 3.0% by weight of or-lerpheny and not less than 0.01% by weight and less than 1.0% by weight of triphenylene based on the total amount of the non-aqueous electrolytic solution are contained.
  - 20. A non-equeous type secondary battery according to claim 17, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohoxylbonzone and not less than 0.01% by weight and less than 1.0% by weight of biphenyl based on the total amount of the non-aqueous electrybic solution are contained.
- 21. A non-aqueous type secondary battery according to claim 17, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohoxyborzene and not less than 0.01% by weight and less than 1.0% by weight of oterphenyl based on the total amount of the non-aqueous electrolytic solution are contained.

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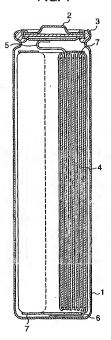
- 22. A non-aqueous type secondary battery according to claim 17, wherein all of not less than 1.0% by weight and not more than 5.0% by weight of cyclohexylbenzene, not less than 0.01% by weight and less than 1.0% by weight of ot-terphenyl and not less than 0.01% by weight of loss than 1.0% by weight of biphenyl based on the lotal amount of the non-equeous electrolytic solution are contained.
- 23. A non-aqueous type secondary bettery according to claim 17, wherein all of o-terphonyl, triphenylene, cyclohex-ylben.zero and biphonyl are contained and the total amount of them is 0.4-5% by weight based on the total amount of the non-aqueous electricity is solution.
- 10 24. A non-equecus type aecondary battery having a positive electrode comprising a material containing a lithium-containing motal order as a positive electrode curve material, a negative electrode comprising a material containing graphite as a negative electrode are material and a non-aqueous electropyle solution, characterized in that he non-aqueous electropyle solution, characterized in the heart of the properties of the properties electropyle configurations a non-aqueous electropyle solution, characterized in the heart of the properties o
- 25. A non-acueous type secondary battery according to claim 24, wherein two or more organic compounds selected from o-terphenyl, tirphenylene, cyclohexylbenzene and biphenyl are contained as the organic compounds differing in oxidative polymerization potential.
  - 26. A non-aqueous type secondary battery according to claim 24, wherein not less than 1.0% by weight and not more than 3.0% by weight of o terphenyl and not less than 0.01% by weight and less than 1.0% by weight of triphenylene based on the total amount of the non-equeous descriptive colution are contained.
  - 27. A non-aqueous type secondary battery according to claim 24, wherein not less than 1.0% by weight and not more than 5.0% by weight of cyclohoxy/benzone and not less than 0.01% by weight and less than 1.0% by weight of biphony/ based on the total amount of the non-aqueous electrolytic solution are contained.
  - 28. A non-equeous type secondary battery according to claim 24, wherein not less than 1.0% by weight and not more than 6.0% by weight of cyclohexylibenziene and not less than 0.01% by weight and less than 1.0% by weight of otephenry based on the total amount of the non-aqueous electrolite solution are contained.
- 5 29. A non-squeous type secondary battery according to claim 24, wherein all of not loss than 1.0% by weight and not more than 5.0% by weight of cyclohexybenzene, not less than 0.01% by weight and less than 1.0% by weight and less than 1.0% by weight of chephenyl and not less than 0.01% by weight and less than 1.0% by weight of biphenyl based on the total amount of the non-aqueous electroytic solution are contained.
- 49 30. A non-aqueous type secondary battery according to claim 24, wherein all of o-terphenyl, triphenylene, cyclohexythenzene and biphenyl are contained and the total amount of them is 0.4-5% by weight based on the total amount of the non-aqueous electrolytic solution.
- 31. A non-equeous type secondary battery according to claim 24, wherein the cyclic carbonate is at least one compound selected from ethylene carbonate (EC), propylene carbonate (PC), butylene carbonate (BC) and vinylene carbonate (BC).
- 32. A non-equeous type secondary battery according to claim 24, wherein the chain carbonate is at least one compound selected from directly carbonate (EMC), dethyl carbonate (DEC), ethylmethyl carbonate (EMC), methylpropyl carbonate (HMC) and ethylpropyl carbonate (EPC).

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# INTERNATIONAL SEARCH REPORT

International application No. PCT/JP01/07434

A. CLASSIFICATION OF SUBJECT MATTER Int.Cl<sup>7</sup> H01/1 10/40

According to International Patent Classification (IPC) or to both national classification and IPC

According to International Patent Classifications (IPC) or to 2021: mannest classification and irr.

B. FIELDS SEARCIED
Milharms documentation searched (classification system followed by classification symbols)
Int. CL7 HOLM 10/40, 6/00-5/22

Documentation searched other than minimum documentation to the cateof that such documents are isolated in the fields searched

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Kokai Jitsuyo Shinan Koho 1971-2001 Jitsuyo Shinan Toroku Koho 1996-2001

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

# C. DOCUMENTS CONSIDERED TO BE RELEVANT Carennys Citation of de-

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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A	JP 2000-58116 A (Samyo Electric Co., Ltd.), 25 February, 2000 (25.02.00), Full text (Family: none)	1-32
A	US 5879834 h (NEC Moil Energy Ltd.,), 09 March, 1999 (09.03.99), Full text & UP 9-106835 h	1-32
PA	JP 2001-210364 A (Mitsubishi Chemical Corporation), 03 August, 2001 (03.08.01), Full text (Family: none)	1-32

# Further documents are listed in the continuation of Rox C See potent families

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